

GenCore version 5.1.6
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OM protein - protein search, using sw model

Run on: June 25, 2003, 14:20:41 ; Search time 31.5 Seconds
(without alignments)
444.169 Million cell updates/sec

Title: US-09-622-613B-6

Perfect score: 583
Sequence: 1 MOEWLTFQKKHLFTNRDVC.....TFCVTCENQAPVHFVGVC 105

Scoring table: BLOSUM62
Gapop 10.0, Gapext 0.5

Searched: 908470 seqs, 133250620 residues

Total number of hits satisfying chosen parameters: 908470

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%
Listing first 45 summaries

Database: A_Geneseq_101002.*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	583	100.0	105	20	AAV28867
2	580	99.5	105	20	AAV28869
3	578	99.1	104	20	AAV28865
4	578	99.1	105	20	AAV28871
5	578	99.1	127	20	AAV28879
6	575	98.6	104	20	AAV28866
7	573	98.3	104	20	AAV28870
8	558	95.7	104	18	AAW06544
9	558	95.7	105	18	AAW35123
10	558	95.7	105	20	AAV39400

11	558	95.7	355	18	AAW35125	R. pipiens recombi
12	558	95.7	358	18	AAW35130	R. pipiens recombi
13	556	95.4	104	18	AAW30301	Recombinant onc pr
14	556	95.4	104	22	AAH31666	Amino acid sequenc
15	556	95.4	112	18	AAW35118	R. pipiens recombi
16	556	95.4	251	18	AAW35134	R. pipiens recombi
17	556	95.4	254	18	AAW35135	R. pipiens recombi
18	556	95.4	355	18	AAW35129	R. pipiens recombi
19	556	95.4	355	18	AAW35133	R. pipiens recombi
20	556	95.4	366	18	AAW35132	R. pipiens recombi
21	556	95.4	379	18	AAW35126	R. pipiens recombi
22	553	94.9	104	12	AAH2344	Protein with activ
23	553	94.9	104	15	AAH47303	ONCONASE (pharmace
24	553	94.9	104	17	AAW0736	Protein derived fr
25	553	94.9	104	18	AAW06543	Antitumor protein
26	553	94.9	104	18	AAW14065	Onconase (RTM) pro
27	553	94.9	104	20	AAH33322	Frog onconase prot
28	553	94.9	104	20	AAH88233	Rana pipiens RNase
29	551	94.5	105	18	AAW35116	R. pipiens recombi
30	551	94.5	106	18	AAW35122	R. pipiens recombi
31	551	94.5	107	18	AAW35117	R. pipiens recombi
32	550	94.3	104	18	AAW30302	Recombinant onc pr
33	550	94.3	105	18	AAW35115	R. pipiens recombi
34	548	94.0	104	18	AAW18224	R. pipiens recombi
35	548	94.0	104	22	AAH31667	Amino acid sequenc
36	547	93.8	358	18	AAW35127	R. pipiens recombi
37	547	93.8	365	18	AAW35131	R. pipiens recombi
38	528	90.6	107	18	AAW35120	R. pipiens recombi
39	495	84.9	360	18	AAW35128	R. pipiens recombi
40	483.5	82.9	111	18	AAW35121	R. pipiens recombi
41	445	76.3	83	18	AAW35119	R. pipiens clone R
42	445	76.3	83	20	AAW88234	Rana pipiens RNase
43	289	49.6	111	20	AAV33321	Frog lectin protel
44	286.5	49.1	111	20	AAV28873	Recombinant Met(-1
45	282.5	48.5	111	20	AAV28876	Recombinant Met(-1

ALIGNMENTS

AAV28867	standard: Protein: 105 AA.
AAV28867	
AC	AAV28867: (first entry)
AC	25-JAN-2000
DT	Recombinant Met(-1) RapLRI.
XX	
DE	Recombinant Met(-1) RapLRI.
XX	
KW	Recombinant Met(-1) Rana pipiens ribonuclease; RapLRI; CD22; RNase;
KW	covalently bound; Ig2 antibody; ligand binding moiety; cancerous B cell;
KW	Kaposi's sarcoma; human chorionic gonadotropin; hCG; signal peptide;
KW	recombinant ribonuclease; cytotoxic fusion protein; cancer; frog;
KW	autoimmune disease.
OS	Rana pipiens.
OS	Synthetic.
XX	
FT	Key
FT	Misc-difference 1
FT	Location/Qualifiers
XX	/note="Met not found in wild type RapLRI"
PN	WO950398-A2.
XX	
PD	07-OCT-1999.
XX	
PF	26-MAR-1999; 99WO-US06641.
XX	
PR	27-MAR-1998; 98US-0079751.
XX	
PA	(USSH) US DEPT HEALTH & HUMAN SERVICES.
XX	

PI	Newton DL, Rybak SM;
DR	WPI: 1999-610847/52.
DR	N-PDB; AAZ08126.
XX	
PT	New recombinant ribonucleases, used for killing target cells, e.g. for treating cancers, viral infections or autoimmune diseases
PS	Claim 34; Page 57; 71pp; English.
XX	
CC	The present sequence is a recombinant Rana pipiens ribonuclease (RapLRI) protein with Met at position 1. Carboxy terminal end of recombinant RapLRI has a covalently bound ligand binding moiety, which can be a LL2 antibody directed against CD22 on cancerous B cells or human chorionic gonadotropin (hCG) effective against Kaposi's sarcoma cells. Recombinant ribonuclease can be expressed in bacteria without an N-terminal methionine due to the presence of a signal peptide that is cleaved by bacteria. The soluble expression of ribonuclease allows the proteins to be fused in-frame with ligand binding moieties to form cytotoxic fusion proteins. They can be used for treatment of cancer and autoimmune diseases.
CC	
SQ	Sequence 105 AA:
QY	Query Match 100.0%; Score 583; DB 20; Length 105; Best Local Similarity 100.0%; Pred. No. 4..2e-63; Matches 105; Conservative 0; Mismatches 0; Indels 0; Gaps 0
Db	1 MODWLTFQKKHLJNTRDVCNNIMSTLFHCKDKNTFTISRPPEVKAICGIIASKNVLT 60 1 MODWLTFQKKHLJNTRDVDCNNIMSGNFHCXDKNTFTISRPEPKAICKGIISKNVLT 60
OY	61 TSEFLLSDCNYTSRPECKYKRLKSKSTVFECVTCENQAPVHFVGVGHG 105 61 TSEFLLSDCNYTSRPECKYKRLKSKSTVFECVTCENQAPVHFVGVGHG 105
Db	
RESULT 2	
AAAY28869	
ID	AAAY28869 standard; Protein: 105 AA.
AC	
XX	AAAY28869;
DT	25-JAN-2000 (first entry)
XX	
DE	Recombinant Met(-1) RapLRI Met23Leu-(His)6 protein.
KW	
RX	Recombinant Met(-1) Rana pipiens ribonuclease Met23Leu-(His)6; RapLRI; CD22; covalently bound; LL2 antibody; ligand binding moiety; RNase; cancerous B cell; Kaposi's sarcoma; human chorionic gonadotropin; hCG; signal peptide; recombinant ribonuclease; cytotoxic fusion protein; cancer; Ilog; autoimmune disease.
XX	
OS	Rana pipiens.
OS	Synthetic.
XX	
FH	Key Location/Qualifiers
FT	Misc-difference 1 /note= "(His)6 histidine tag attached to N-terminal Met"
FT	Misc-difference 1 /note= "Met not found in wild type RapLRI"
FT	Misc-difference 24 /note= "Wild type Met replaced with Leu"
XX	
PN	MO950398-AA2.
PD	07-OCT-1999.
XX	
PF	26-MAR-1999; 99MO-US06641.
XX	
PR	27-MAR-1998; 98US-0079751.
XX	
VA	(USSH) US DEPT HEALTH & HUMAN SERVICES.

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XX XX Newton DL, Rybak SM;
XX PI
XX WP1; 1999-610847/52.
DR N-PSDB; AA208127.
XX
XX New recombinant ribonucleases, used for killing target cells, e.g. for
XX PT treating cancers, viral infections or autoimmune diseases
XX PS
XX Claim 4; Page 59; 71pp; English.
XX
XX The present sequence is a recombinant Rana pipiens ribonuclease protein
CC (RapiR1) with Met at position 1 attached to (His)6 tag and Met24Leu.
CC Carboxy terminal end of recombinant RapiR1 has a covalently bound ligand
CC binding moiety, which can be a LL2 antibody directed against CD22 on
CC cancerous B cells or human chorionic gonadotropin (hCG) effective
CC against Kaposi's sarcoma cells. Recombinant ribonucleases can be
CC expressed in bacteria without an N-terminal methionine due to the
CC presence of a signal peptide that is cleaved by bacteria. The soluble
CC expression of ribonuclease allows the proteins to be fused in-frame with
CC ligand binding moieties to form cytotoxic fusion proteins. They can be
CC used for treatment of cancer and autoimmune diseases.
XX
XX
SQ Sequence 105 AA;
XX
XX
XX Query Match 99.5%; Score 580; DB 20; Length 105;
XX Best Local Similarity 99.0%; Pred. No. 9,7e-63;
XX Matches 104; Conservative 1; Mismatches 0; Indels 0; Gaps 0
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XX QY 1 MODMLTFQKKHLTNFQDVCCNNIMSTNLFHCKDKNTFTYSRPEPKAICKGIIASKNVL 60
XX |||||||
XX 1 MODMLTFQKKHLTNFQDVCCNNILSTNLFHCKDKNTFTYSRPEPKAICKGIIASKNVL 60
XX
XX QY 1 TSEFYLSDCNVTSRPCKYKLRKSTNFTCVTCENQAPVHFVGVGHC 105
XX |||||||
XX 1 TSEFYLSDCNVTSRPCKYKLRKSTNFTCVTCENQAPVHFVGVGHC 105
XX
XX Db
XX 61 TSEFYLSDCNVTSRPCKYKLRKSTNFTCVTCENQAPVHFVGVGHC 105
XX
XX
XX RESULT 3
XX AA128865
XX ID AAY28865 standard; Protein; 104 AA.
XX AC
XX AAY28865;
XX
XX DT 25-JAN-2000 (first entry)
XX
XX DE Rana pipiens liver ribonuclease (RapiR1).
XX
XX Rana pipiens liver ribonuclease; RapiR1; covalently bound; LL2 antibody;
KW ligand binding moiety; CD22; cancerous B cell; Kaposi's Sarcoma; frog;
KW human chorionic gonadotropin; hCG; recombinant ribonuclease; RNase;
KW signal peptide; cytotoxic fusion protein; cancer; autoimmune disease.
XX
XX Rana pipiens.
XX
XX OS
XX
XX WO9503398-A2.
XX PN
XX PD 07-OCT-1999.
XX
XX PF 26-MAR-1999; 99WO-US06641.
XX
XX PR 27-MAR-1998; 98US-0079751.
XX
XX PA (USSH ) US DEPT HEALTH & HUMAN SERVICES.
XX
XX PI Newton DL, Rybak SM;
XX
XX WP1; 1999-610847/52.
XX DR N-PSDB; AA208124.
XX
XX New recombinant ribonucleases, used for killing target cells, e.g. for
XX PT treating cancers, viral infections or autoimmune diseases
XX

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PS Claim 1; Page 55; 71pp; English.

XX The present sequence is Rana pipiens liver ribonuclease (RapLr1)

CC protein. Carboxy terminal end of RapLr1 has a covalently bound

CC ligand binding moiety, which can be a LL2 antibody directed against

CC CD22 on cancerous B cells or human chorionic gonadotropin (hCG)

CC effective against Kaposi's Sarcoma cells. Recombinant ribonucleases can

CC be expressed in bacteria without an N-terminal methionine due to the

CC presence of a signal peptide that is cleaved by bacteria. The soluble

CC expression of ribonuclease allows the proteins to be fused in-frame with

CC ligand binding moieties to form cytotoxic fusion proteins. They can be

CC used for treatment of cancer and autoimmune diseases.

XX

SQ Sequence 104 AA;

Query Match 99.1%; Score 578; DB 20; Length 104;

Best Local Similarity 100.0%; Pred. No. 1.7e-62;

Matches 104; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 QDWLTFQKKHLNTRDVCNNIMSTNLFHCKDKNTFTYSRPPVKAICKGIASKNVLT 61

DB 1 QDWLTFQKKHLNTRDVCNNIMSTNLFHCKDKNTFTYSRPPVKAICKGIASKNVLT 60

QY 62 SEFYLSDCNVTSRPCKYKLLKSTNTPFCVTCENQAPVHFVGHC 105

DB 61 SEFYLSDCNVTSRPCKYKLLKSTNTPFCVTCENQAPVHFVGHC 104

RESULT 4

AAI28871

ID AAY28871 standard; Protein; 105 AA.

XX

AC AAY28871;

XX

DT 25-JAN-2000 (first entry)

DE Recombinant Met(-1) RapLr1 GlnSer amino acid sequence.

XX

XX Recombinant Met(-1) Rana pipiens ribonuclease GlnSer; RapLr1; CD22;

KM covalently bound; LL2 antibody; ligand binding moiety; cancerous B cell;

KM Kaposi's sarcoma; human chorionic gonadotropin; hCG; signal peptide;

KM recombinant ribonuclease; cytotoxic fusion protein; cancer; frog;

KW autoimmune disease; RNase.

XX

OS Rana pipiens.

XX Synthetic.

OS

FH Key Location/Qualifiers

FT MISC-difference 1 /note= "Met not found in wild type RapLr1"

FT MISC-difference 2 /note= "Wild type Gln replaced with Ser"

FT

XX

PN WO950398-A2.

XX

XX 07-OCT-1999.

PD

XX

XX 26-MAR-1999; 99WO-US06641.

PF

XX

XX 27-MAR-1998; 98US-0079751.

PR

XX

XX (USSH) US DEPT HEALTH & HUMAN SERVICES.

PA

XX

XX Newton DL, Rybak SM;

PI

XX

DR WPI, 1999-610847/52.

DR

XX

XX N-PSDB; AA208129.

DR

XX

PT New recombinant ribonucleases, used for killing target cells, e.g. for

PT treating cancers, viral infections or autoimmune diseases

XX

PS Claim 34; Page 61; 71pp; English.

XX

CC The present sequence is a recombinant Rana pipiens ribonuclease (RapLr1)

CC protein with Met at position 1 and Gln2Ser. Carboxy terminal end of

CC recombinant RapLr1 has a covalently bound ligand binding moiety, which

CC can be a LL2 antibody directed against CD22 on cancerous B cells or human

CC chorionic gonadotropin (hCG) effective against Kaposi's sarcoma cells.

CC Recombinant ribonucleases can be expressed in bacteria without an N-

CC terminal methionine due to the presence of a signal peptide that is

CC cleaved by bacteria. The soluble expression of ribonuclease allows the

CC proteins to be fused in-frame with ligand binding moieties to form

CC cytotoxic fusion proteins. They can be used for treatment of cancer and

CC autoimmune diseases.

XX

SQ Sequence 105 AA;

Query Match 99.1%; Score 578; DB 20; Length 105;

Best Local Similarity 99.0%; Pred. No. 1.7e-62;

Matches 104; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

QY 1 QDWLTFQKKHLNTRDVCNNIMSTNLFHCKDKNTFTYSRPPVKAICKGIASKNVLT 60

DB 1 MSDWLTFQKKHLNTRDVCNNIMSTNLFHCKDKNTFTYSRPPVKAICKGIASKNVLT 60

QY 61 TSEFYLSDCNVTSRPCKYKLLKSTNTPFCVTCENQAPVHFVGHC 105

DB 61 TSEFYLSDCNVTSRPCKYKLLKSTNTPFCVTCENQAPVHFVGHC 105

RESULT 5

AAI28879

ID AAY28879 standard; Protein; 127 AA.

XX

XX

AC AAY28879;

XX

DT 25-JAN-2000 (first entry)

DE Rana pipiens Clone 5a1b ribonuclease.

XX

XX Rana pipiens ribonuclease Clone 5a1b; RapLr1; covalently bound; RNase;

KM LL2 antibody; ligand binding moiety; CD22; cancerous B cell; oncogene;

KM Kaposi's Sarcoma; human chorionic gonadotropin; hCG; cancer;

KM recombinant ribonuclease; frog; signal peptide; cytotoxic fusion protein;

KW autoimmune disease.

XX

OS Rana pipiens.

XX

FH Key Location/Qualifiers

FT Peptide 1..23

FT /label= "Signal-peptide"

FT /note= "Putative"

FT 24..127

FT Protein /label= Rana-pipiens_Clone_5a1b_ribonuclease

FT

XX

PN WO950398-A2.

XX

XX 07-OCT-1999.

PD

XX

XX 26-MAR-1999; 99WO-US06641.

PF

XX

XX 27-MAR-1998; 98US-0079751.

PR

XX

XX (USSH) US DEPT HEALTH & HUMAN SERVICES.

PA

XX

XX Newton DL, Rybak SM;

PI

XX

DR WPI, 1999-610847/52.

DR

XX

XX N-PSDB; AA208136.

DR

XX

PT New recombinant ribonucleases, used for killing target cells, e.g. for

PT treating cancers, viral infections or autoimmune diseases

XX

PS Disclosure; Page 69; 71pp; English.

XX

CC The present sequence is a Rana pipiens Clone 5a1b ribonuclease (RapLr1).

CC It is encoded by Clone 5a1b cDNA obtained from Rana pipiens liver mRNA
 CC library. It exhibits differences with Onconase (RPM) at amino acid
 CC residues 11, 20, 85 and 103. Carboxy terminal end of RapLRI has a
 CC covalently bound ligand binding moiety, which can be a Lf2 antibody
 CC directed against CD22 on cancerous B cells or human chorionic
 CC gonadotropin (hCG) effective against Kaposi's Sarcoma cells. Recombinant
 CC ribonucleases can be expressed in bacteria without an N-terminal
 CC methionine due to the presence of a signal peptide that is cleaved by
 CC bacteria. The soluble expression of ribonuclease allows the proteins to
 CC be fused in-frame with ligand binding moieties to form cytotoxic fusion
 CC proteins. They can be used for treatment of cancer and autoimmune
 CC diseases.

XX Sequence 127 AA;

Query Match 99.1%; Score 578; DB 20; Length 127;
 Best Local Similarity 100.0%; Pred. No. 2.2e-62;
 Matches 104; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2 QDWLTFQKHLLTNRDVCNNIMSTNLFHCKDKNTFTYSRPEPYKAICKGIASKNVLT 61
 |||||||
 Db 24 QDWLTFQKHLLTNRDVCNNIMSTNLFHCKDKNTFTYSRPEPYKAICKGIASKNVLT 83

OY 62 SEFTLSDCNVTSRPCKTKLKSTNFCVTCENQAPVHFVGHC 105
 |||||||
 Db 84 SEFTLSDCNVTSRPCKTKLKSTNFCVTCENQAPVHFVGHC 127

RESULT 6

AAV28866
 ID AAV28866 standard; Protein: 104 AA.

AC AAV28866;

XX 25-JAN-2000 (first entry)

DE Recombinant RapLRI Met23Leu amino acid sequence.

KW Recombinant Rana pipiens ribonuclease; RapLRI Met23Leu; covalently bound;
 KM Lf2 antibody; ligand binding moiety; CD22; cancerous B cell; RNase;
 KM Kaposi's sarcoma; human chorionic gonadotropin; hCG; signal peptide;
 KW recombinant ribonuclease; cytotoxic fusion protein; cancer; frog;
 KW autoimmune disease.

XX Rana pipiens.
 OS Synthetic.

XX Key Location/Qualifiers

FT Misc-difference 23 /note= "Wild type Met replaced with Leu"

XX MO9950398-A2.

XX 07-OCT-1999.

XX 26-MAR-1999; 99WO-US06641.

XX 27-MAR-1998; 98US-0079751.

XX (USSH) US DEPT HEALTH & HUMAN SERVICES.

XX Newton DL, Rybak SM;

XX WPI; 1999-610847/52.

XX N-PSDB; AA208125.

PT New recombinant ribonucleases, used for killing target cells, e.g. for
 PT treating cancers, viral infections or autoimmune diseases
 XX Claim 34; Page 56; 71pp; English.

CC The present sequence is a recombinant Rana pipiens ribonuclease (RapLRI)
 CC protein with Met23Leu. Carboxy terminal end of recombinant RapLRI has a

CC covalently bound ligand binding moiety, which can be a Lf2 antibody
 CC directed against CD22 on cancerous B cells or human chorionic
 CC gonadotropin (hCG) effective against Kaposi's sarcoma cells. Recombinant
 CC ribonucleases can be expressed in bacteria without an N-terminal
 CC methionine due to the presence of a signal peptide that is cleaved by
 CC bacteria. The soluble expression of ribonuclease allows the proteins to
 CC be fused in-frame with ligand binding moieties to form cytotoxic fusion
 CC proteins. They can be used for treatment of cancer and autoimmune
 CC diseases.

XX Sequence 104 AA;

Query Match 98.6%; Score 575; DB 20; Length 104;
 Best Local Similarity 99.0%; Pred. No. 3.9e-62;
 Matches 103; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

OY 2 QDWLTFQKHLLTNRDVCNNIMSTNLFHCKDKNTFTYSRPEPYKAICKGIASKNVLT 61
 |||||||
 Db 1 QDWLTFQKHLLTNRDVCNNIMSTNLFHCKDKNTFTYSRPEPYKAICKGIASKNVLT 60

OY 62 SEFTLSDCNVTSRPCKTKLKSTNFCVTCENQAPVHFVGHC 105
 |||||||
 Db 61 SEFTLSDCNVTSRPCKTKLKSTNFCVTCENQAPVHFVGHC 104

RESULT 7

AAV28870
 ID AAV28870 standard; Protein: 104 AA.

AC AAV28870;

XX 25-JAN-2000 (first entry)

DE Recombinant RapLRI GlnSer amino acid sequence.

KW Recombinant Rana pipiens ribonuclease; RapLRI GlnSer; covalently bound;
 KM Lf2 antibody; ligand binding moiety; CD22; cancerous B cell; frog;
 KM Kaposi's sarcoma; human chorionic gonadotropin; hCG; signal peptide;
 KW recombinant ribonuclease; cytotoxic fusion protein; cancer; RNase;
 KW autoimmune disease.

XX Rana pipiens.
 OS Synthetic.

XX Key Location/Qualifiers

FT Misc-difference 1 /note= "Wild type Gln replaced with Ser"

XX MO9950398-A2.

XX 07-OCT-1999.

XX 26-MAR-1999; 99WO-US06641.

XX 27-MAR-1998; 98US-0079751.

XX (USSH) US DEPT HEALTH & HUMAN SERVICES.

XX Newton DL, Rybak SM;

XX WPI; 1999-610847/52.

XX N-PSDB; AA208128.

PT New recombinant ribonucleases, used for killing target cells, e.g. for
 PT treating cancers, viral infections or autoimmune diseases
 XX Claim 34; Page 60; 71pp; English.

CC The present sequence is a recombinant Rana pipiens ribonuclease (RapLRI)
 CC protein with GlnSer. Carboxy terminal end of recombinant RapLRI has a
 CC covalently bound ligand binding moiety, which can be a Lf2 antibody
 CC directed against CD22 on cancerous B cells or human chorionic
 CC gonadotropin (hCG) effective against Kaposi's sarcoma cells. Recombinant

CC ribonucleases can be expressed in bacteria without an N-terminal methionine due to the presence of a signal peptide that is cleaved by CC bacteria. The soluble expression of ribonuclease allows the proteins to be fused in-frame with ligand binding moieties to form cytotoxic fusion CC proteins. They can be used for treatment of cancer and autoimmune diseases.

XX Sequence 104 AA;

Query Match 98.3%; Score 573; DB 20; Length 104;
Best Local Similarity 100.0%; Pred. No. 6.9e-62;
Matches 103; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 DMLTFQKKHLTNTRDVCNNIMSTNLFHCKDKNTFYISRPPEVKAICKGIISKNVLTTS 62
|||||
DB 2 DMLTFQKKHLTNTRDVCNNIMSTNLFHCKDKNTFYISRPPEVKAICKGIISKNVLTTS 61

QY 63 EFYLSDCNVTSRPCKYKLLKSKTNFCVTCENQAPVHFGVGHG 105
|||||
DB 62 EFYLSDCNVTSRPCKYKLLKSKTNFCVTCENQAPVHFGVGHG 104

RESULT 8
AAW06544
ID AAW06544 standard; Protein: 104 AA.

XX AAW06544;

DT 22-AUG-1997 (first entry)

DE Antitumour protein from Rana pipiens oocytes.

KW Tumour; chemotherapy; radiotherapy; frog.

OS Rana pipiens.

PN W09639428-A1.

PD 12-DEC-1996.

PF 03-JUN-1996; 96WO-US08304.

PR 06-JUN-1995; 95US-0467955.

PA (ALFA-) ALFACELL CORP.

PI Ardelt WJ;

DR WPI: 1997-043063/04.

XX Antitumour proteins from Rana pipiens oocyte(s) - have fewer
PT disadvantages than chemotherapy, surgery and radiotherapy
PS

PS Claim 8; Page 28; 45pp; English.

XX The present sequence is a specifically claimed example of an
CC antitumour protein from the genetic protein in AAW18224, with the
CC molecular weight 12000. This is one of two preferred proteins (the
CC other in AAW06543) that have been isolated from Rana pipiens oocytes.
CC Both proteins have a blocked amino terminal group and are essentially
CC free of carbohydrates. The proteins are used to treat tumours. Use of
CC the peptides has fewer disadvantages than chemotherapy, radiotherapy
CC and surgery in the treatment of tumours.

XX Sequence 104 AA;

Query Match 95.7%; Score 558; DB 18; Length 104;
Best Local Similarity 96.2%; Pred. No. 4.6e-60;
Matches 100; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

QY 2 QDMLTFQKKHLTNTRDVCNNIMSTNLFHCKDKNTFYISRPPEVKAICKGIISKNVLT 61
:|||||
DB 1 EDMLTFQKKHLTNTRDVCNNIMSTNLFHCKDKNTFYISRPPEVKAICKGIISKNVLT 60

QY 62 SEFYLSDCNVTSRPCKYKLLKSKTNFCVTCENQAPVHFGVGHG 105
|||||
DB 61 SEFYLSDCNVTSRPCKYKLLKSKTNFCVTCENQAPVHFGVGHG 104

RESULT 9
AAW35123
ID AAW35123 standard; Protein: 105 AA.

XX AAW35123;

DT 20-APR-1998 (first entry)

DE R. pipiens recombinant RNase protein [Met-(1)]rnc.

KW RNase A; ribonuclease; cytotoxic; onconase; nOnc; immunofusion;
tumor cell growth; frog.

OS Rana pipiens.

PN W09731116-A2.

PD 28-AUG-1997.

PF 19-FEB-1997; 97WO-US02588.

PR 21-FEB-1996; 96US-0011800.

PA (USSH) US DEPT HEALTH & HUMAN SERVICES.

PI Boque L, Newton DL, Rybak SM, Wlodawer A;

DR WPI: 1997-435168/40.

DR N-PSDB; AAT94959.

PT Ribonuclease molecules based on native Onconase - used for killing
cells, particularly tumour cells

PS Disclosure; Pages 65-66; 90pp; English.

XX AAW35115 to AAW35123 encode recombinant proteins (rnc) which are
CC modifications of the RNase Onconase (RNM) (rnc). Such novel
CC ribonuclease molecules are highly cytotoxic and can be used alone or to
CC form chemical conjugates or to target recombinant immunofusions. They are
CC used particularly for decreasing tumour cell growth. They can also be
CC used for cell separation in vitro by selectively killing unwanted types
CC of cells, e.g. in bone marrow prior to transplantation into a patient
CC undergoing marrow ablation by radiation, or for killing leukaemia cells
CC or T-cells that would cause graft versus host disease. The toxins can
CC also be used to selectively kill unwanted cells in culture. The new
CC ribonucleases have increased cytotoxic activity compared to rnc and also
CC lower immunogenicity in humans.

XX Sequence 105 AA;

Query Match 95.7%; Score 558; DB 18; Length 105;
Best Local Similarity 95.2%; Pred. No. 4.6e-60;
Matches 100; Conservative 3; Mismatches 2; Indels 0; Gaps 0;

QY 1 MODLTFQKKHLTNTRDVCNNIMSTNLFHCKDKNTFYISRPPEVKAICKGIISKNVLT 60
:|||||
DB 1 MEDMLTFQKKHLTNTRDVCNNIMSTNLFHCKDKNTFYISRPPEVKAICKGIISKNVLT 60

QY 61 TSEFYLSDCNVTSRPCKYKLLKSKTNFCVTCENQAPVHFGVGHG 105
|||||
DB 61 TSEFYLSDCNVTSRPCKYKLLKSKTNFCVTCENQAPVHFGVGHG 105

RESULT 10
AAV39400
ID AAV39400 standard; Protein: 105 AA.

XX

```

AC AAY39400;
XX
DT 01-DEC-1999 (first entry)
XX
DE Recombinant frog Onconase.
XX
KW Ribonuclease; protein synthesis; inhibition; cancer; cytotoxic.
XX
OS Rana pipiens.
XX
PN W09946389-A1.
XX
PD 16-SEP-1999.
XX
PF 11-MAR-1999; 99WO-US04252.
XX
PR 11-MAR-1998; 98US-0077557.
XX
PA (IMMU-) IMMUNOMEDICS INC.
XX
PI Goldenberg DW, Hansen H, Leung S;
XX
DR MPI: 1999-551416/46.
XX
DR N-PSDB: AAZ19767.
XX
PT
PS
XX
XX Example 1: Fig 1, 42pp; English.
XX
CC This sequence represents recombinant frog onconase. Onconase has
CC ribonuclease and anti-tumour activity. The cDNA was produced via PCR
CC (using primers AAZ19768-Z19769) of two synthetic DNAs whose sequences
CC encoded most of the N-terminal or the C-terminal amino acids of mature
CC onconase. The two PCR products generated encoded either the N-terminal
CC 54 amino acids (minus the initial methionine) or the C-terminal 51 amino
CC acids, and were ligated in frame at an NruI site. The cDNA was then
CC subcloned into a vector e.g., pBluescript, where the ATG initiation
CC codon was ligated to the cDNA. After expression in E. coli, the
CC recombinant protein was purified. The initial N-formyl methionine was
CC cleaved off and the now N-terminal glutamate residue cyclised to form an
CC N-terminal pyroglutamate. The pyroglutamate residue forms part of the
CC phosphate binding pocket of onconase and is essential for both
CC ribonuclease and anti-tumour activity. Onconase is a 12 kD ribonuclease
CC which causes cell death as a result of potent inhibition of protein
CC synthesis by a mechanism involving inactivation of cellular RNA. It is
CC not inhibited by mammalian placental ribonuclease inhibitor, which may
CC explain its enhanced cytotoxicity relative to mammalian enzymes. It has
CC anti-tumour activity against a variety of solid tumours e.g., colon or
CC pancreatic cancers, and can be used alone or in combination with other
CC anti-cancer agents such as tamoxifen. When used as an anti-tumour agent,
CC onconase can be conjugated to a marker which targets it to a specific
CC cell type.
XX
SQ Sequence 105 AA;
XX
OY Query Match 95.7%; Score 558; DB 20; Length 105;
Db Best Local Similarity 95.2%; Pred. No. 4.6e-60;
Matches 100; Conservative 3; Mismatches 2; Indels 0; Gaps 0
OY 1 MODWLTFQKKHLTNRDVDCNNINSTNLFHCKDKNTFTYSRPEPKAICKGIASKNVLT 60
Db 1 MODWLTFOKKHITMTKDDVCNNINSTNLFHCKDKNTFTYSRPEPKAICKGIASKNVLT 60
OY 61 TSEFYLSDCNVTSPRCRYKKLKKSTNTPCVTCENQAPVHFVGCGHC 105
Db 61 TSEFYLSDCNVTSPRCRYKKLKKSTNTPCVTCENQAPVHFVGCGSC 105
ID AAM35125 standard; Protein: 355 AA.
NC AAM35125;

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XX	20-APR-1998	(first entry)
DT	R. pipiens recombinant RNase ronc fusion protein 1.	
XX		
DE	RNase A; ribonuclease; cytotoxic; onconase; nonc; immunofusion;	
XX	tumour cell growth; frog.	
KW		
XX	Rana pipiens.	
OS	Synthetic.	
PN	MO973116-A2.	
XX		
PD	28-AUG-1997.	
XX		
PF	19-FEB-1997; 97WO-US02588.	
XX		
PR	21-FEB-1996; 96US-0011800.	
XX		
PA	(USSH) US DEPT HEALTH & HUMAN SERVICES.	
XX		
PI	Boque L, Newton DL, Rybak SM, Wlodawer A;	
XX		
DR	WPI; 1997-435168/40.	
XX	N-PSDB; AAT94963.	
XX		
PT	Ribonuclease molecules based on native Onconase - used for killing	
XX	cells, particularly tumour cells	
PS	Disclosure: Page 67; 90pp; English.	
XX		
CC	Sequences AAW35125 to AAW35135 represent recombinant fusion proteins	
CC	(ronc) which are modifications of the RNase Onconase (RTM) (nonc). Such	
CC	novel ribonuclease molecules are highly cytotoxic and can be used alone	
CC	or to form chemical conjugates or to target recombinant immunofusions.	
CC	They are used particularly for decreasing tumour cell growth. They can	
CC	also be used for cell separation in vitro by selectively killing unwanted	
CC	types of cells, e.g. in bone marrow prior to transplantation into a	
CC	patient undergoing marrow ablation by radiation, or for killing leukaemia	
CC	cells or T-cells that would cause graft versus host disease. The toxins	
CC	can also be used to selectively kill unwanted cells in culture. The new	
CC	ribonucleases have increased cytotoxic activity compared to nonc and	
XX	also lower immunogenicity in humans.	
SQ	Sequence 355 AA;	
Query Match	95.7%; Score 558; DB 18; Length 355;	
Best Local Similarity	95.2%; Pred. No. 2.2e-59;	
Matches 100; Conservative	3; Mismatches 2; Indels 0; Gaps 0	
QY	1 MODLTFOKKHLTNTRDVDCNNINSTNLFHCKDKKNTFTYSPREPKAICKGIASKNVLT 60	
DB	251 MEDLLTFQKKHITNTRDVDCNINSTNLFHCKDKKNTFTYSPREPKAICKGIASKNVLT 310	
QY	61 TSEFTLSDCANTSRPCKYKRLKKSINTFCVTCENQAPVHFVGVGHC 105	
DB	311 TSEFTLSDCANTSRPCKYKRLKKSINTFCVTCENQAPVHFVGVGSC 355	
RESULT 12		
AAW35130		
ID	AAW35130 standard; Protein; 358 AA.	
XX		
AC	AAW35130;	
XX		
DT	20-APR-1998 (first entry)	
XX		
DE	R. pipiens recombinant RNase ronc fusion protein 6.	
XX		
KW	RNase A; ribonuclease; cytotoxic; onconase; nonc; immunofusion;	
KW	tumour cell growth; frog.	
XX		
OS	Rana pipiens	

OS Synthetic.
 XX
 PN MO9731116-A2.
 XX
 PD 28-AUG-1997.
 XX
 PF 19-FEB-1997; 97MO-US02588.
 XX
 PR 21-FEB-1996; 96US-0011800.
 XX
 PA (USSH) US DEPT HEALTH & HUMAN SERVICES.
 PI Boque L, Newton DL, Rybak SM, Wlodawer A;
 XX
 DR WPI: 1997-435168/40.
 DR N-PSDB; AAT94968.
 XX
 PT Ribonuclease molecules based on native Oncanase - used for killing
 PT cells, particularly tumour cells
 XX
 PS Disclosure; Page 72; 90pp; English.
 XX
 CC Sequences AAM35125 to AAM35135 represent recombinant fusion proteins
 CC (Ponc) which are modifications of the RNase Oncanase (RTM) (nnc). Such
 CC novel ribonuclease molecules are highly cytotoxic and can be used alone
 CC or to form chemical conjugates or to target recombinant immunofusions.
 CC They are used particularly for decreasing tumour cell growth. They can
 CC also be used for cell separation in vitro by selectively killing unwanted
 CC types of cells, e.g. in bone marrow prior to transplantation into a
 CC patient undergoing marrow ablation by radiation, or for killing leukemia
 CC cells or T-cells that would cause graft versus host disease. The toxins
 CC can also be used to selectively kill unwanted cells in culture. The new
 CC ribonucleases have increased cytotoxic activity compared to nnc and
 CC also lower immunogenicity in humans.
 XX
 SO Sequence 358 AA;
 Query Match 95.7%; Score 558; DB 18; Length 358;
 Best Local Similarity 95.2%; Pred. No. 2.3e-59;
 Matches 100; Conservative 3; Mismatches 2; Indels 0; Gaps 0;
 OY 1 MOWLTFQKKHLNTRVDCCNNIMSTNLFHCKDKNTFYSPREPKAICGIIASKNVLT 60
 DB 1 MEDMLTFQKKHINTNRVDCCNNIMSTNLFHCKDKNTFYSPREPKAICGIIASKNVLT 60
 OY 61 TSEFYISDCNVTSPCKYKLTGSTNTPFCVTCENQAPVHFVGVC 105
 DB 61 TSEFYISDCNVTSPCKYKLTGSTNTPFCVTCENQAPVHFVGVC 105
 RESULT 13
 AAM30301
 ID AAM30301 standard; protein; 104 AA.
 XX
 AC AAM30301;
 XX
 DT 09-JUN-1998 (first entry)
 XX
 DE Recombinant onc protein.
 XX
 KW Onc; oncanase; ribonuclease; frog; antitumour; pancreatic cancer;
 KW human immunodeficiency virus type-1; HIV1; replication.
 XX
 OS Rana pipiens.
 OS
 PN MO9738112-A1.
 PN
 PD 16-OCT-1997;
 PD
 PF 04-APR-1997; 97MO-US05675.
 PF
 PR 04-APR-1996; 96US-0626288.
 PR
 XX

PA (USSH) US DEPT HEALTH & HUMAN SERVICES.
 PI Adelt W, Boix E, Vasandani VM, Wu YN, Youle RJ;
 XX
 DR WPI: 1997-512725/47.
 DR
 XX
 PT Recombinant Onc protein with glutamine residue at position 1 -
 PT useful as antitumour and antiviral agent, also as cell culture
 PT selection agent
 XX
 PS Claim 1; Page 28; 35pp; English.
 XX
 CC This sequence represents a recombinant Onc protein comprising a 104 amino-
 CC acid sequence having Gln at position 1. Onc, a ribonuclease from Rana
 CC pipiens oocytes, is known as an antitumour agent (e.g. for treating
 CC pancreatic cancer) and inhibitor of human immunodeficiency virus type-1
 CC replication. It can be used therapeutically or as a cell-culture
 CC selection agent, e.g. to identify gene therapy compositions able to
 CC inhibit tumour growth.
 XX
 SO Sequence 104 AA;
 Query Match 95.4%; Score 556; DB 18; Length 104;
 Best Local Similarity 96.2%; Pred. No. 8e-60;
 Matches 100; Conservative 2; Mismatches 2; Indels 0; Gaps 0;
 OY 2 QDWLTFQKKHLNTRVDCCNNIMSTNLFHCKDKNTFYSPREPKAICGIIASKNVLT 61
 DB 1 QDWLTFQKKHINTNRVDCCNNIMSTNLFHCKDKNTFYSPREPKAICGIIASKNVLT 60
 OY 62 SEFYISDCNVTSPCKYKLTGSTNTPFCVTCENQAPVHFVGVC 105
 DB 61 SEFYISDCNVTSPCKYKLTGSTNTPFCVTCENQAPVHFVGVC 104
 RESULT 14
 AAB31666
 ID AAB31666 standard; protein; 104 AA.
 XX
 AC AAB31666;
 XX
 DT 30-APR-2001 (first entry)
 XX
 DE Amino acid sequence of a frog ribonuclease protein.
 XX
 KW Frog; ribonuclease; ranpirinase; RNase.
 XX
 OS Rana pipiens.
 OS
 FT Key Location/Qualifiers
 FT Modified-site 1 /note="this Gln is autocyclised to pyroglutamic acid"
 FT
 XX
 PN US6175003-B1.
 PN
 PD 16-JAN-2001.
 PD
 PF 10-SEP-1999; 99US-0394268.
 PF
 PR 10-SEP-1999; 99US-0394268.
 PR
 PA (ALFA-) ALFACELL CORP.
 PA
 PI Saxena SK;
 PI
 DR WPI: 2001-167808/17.
 DR
 PT New nucleic acids encoding a ribonuclease (Rnase), useful for the
 PT precise targeting of Rnase to a predetermined cell receptor -
 PT
 PS Claim 1; Columns 5-6; 7pp; English.
 PS
 CC The present sequence represents a frog ribonuclease protein (ranpirinase)

CC (RNase). The specification describes a synthetic ribonuclease protein,
CC in which the addition of cysteine in the ribonuclease facilitates the
CC chemical linking of a targeting molecule by the single reactive
CC sulfhydryl group. The specification also describes a method for the
CC production of rnapinase using DNA technology instead of processing
CC biological material. The re-engineering of the protein molecule allows
CC easier attachment to a targeting molecule thereby making it possible for
CC the ribonuclease to be delivered to a particular cell receptor where it
CC might be most effective.
XX

SO Sequence 104 AA;

Query Match 95.4%; Score 556; DB 22; Length 104;
Best Local Similarity 96.2%; Pred. No. 8e-60;
Matches 100; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

OY 2 QDWLTFQKKHLTNRDVDCNNIMSTNLFHCKDKNFTIYSRPEPKAICKGIASKNVLT 61
DB 1 QDWLTFQKKHLTNRDVDCNNIMSTNLFHCKDKNFTIYSRPEPKAICKGIASKNVLT 60

OY 62 SEFYLSDCNVTSRPCCKYKLLKSTNFCVTCENQAPVHFGVGC 105
DB 61 SEFYLSDCNVTSRPCCKYKLLKSTNFCVTCENQAPVHFGVGC 104

RESULT 15

AAW35118
ID AAW35118 standard; Protein: 112 AA.

AC AAW35118;

DT 20-APR-1998 (first entry)

DE R. p1piens recombinant RNase protein NLSmetSerrOnc.

KM RNase A: ribonuclease; cytotoxic; onconase; nonc; immunofusion;
KW tumour cell growth; frog.

OS Rana pipiens.

PN WO9731116-A2.

PD 28-AUG-1997.

PE 19-FEB-1997; 97WO-US02588.

PR 21-FEB-1996; 96US-0011800.

PA (USSH) US DEPT HEALTH & HUMAN SERVICES.

PI Boque L, Newton DL, Rybak SM, Wlodawer A;

DR WPI; 1997-435168/40.

DR N-PSDB; AAT94955.

PT Ribonuclease molecules based on native Onconase - used for killing

PS cells, particularly tumour cells

PS Claim 18; Page 63; 90pp; English.

CC AAW35115 to AAW35123 encode recombinant proteins (rOnc) which are
CC modifications of the RNase Onconase (rOnc). Such novel
CC ribonuclease molecules are highly cytotoxic and can be used alone or to
CC form chemical conjugates or to target recombinant immunofusions. They are
CC used particularly for decreasing tumour cell growth. They can also be
CC used for cell separation in vitro by selectively killing unwanted types
CC of cells, e.g. in bone marrow prior to transplantation into a patient
CC undergoing marrow ablation by radiation, or for killing leukaemia cells
CC or T-cells that would cause graft versus host disease. The toxins can
CC also be used to selectively kill unwanted cells in culture. The new
CC ribonucleases have increased cytotoxic activity compared to nOnc and also
CC lower immunogenicity in humans.
XX

SO Sequence 112 AA;

Query Match 95.4%; Score 556; DB 18; Length 112;
Best Local Similarity 95.2%; Pred. No. 8.9e-60;
Matches 100; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

OY 1 MODWLTFQKKHLTNRDVDCNNIMSTNLFHCKDKNFTIYSRPEPKAICKGIASKNVLT 60
DB 8 MSDWLTFQKKHLTNRDVDCNNIMSTNLFHCKDKNFTIYSRPEPKAICKGIASKNVLT 67

OY 61 TSEFYLSDCNVTSRPCCKYKLLKSTNFCVTCENQAPVHFGVGC 105
DB 68 TSEFYLSDCNVTSRPCCKYKLLKSTNFCVTCENQAPVHFGVGC 112

Search completed: June 25, 2003, 14:48:38
Job time : 32.5 secs